=> d 11 L1 HAS NO ANSWERS L1 STR

VPA 24-9/10/11 U NODE ATTRIBUTES: NSPEC IS R AT 21 DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RSPEC 10 4 NUMBER OF NODES IS 26

STEREO ATTRIBUTES: NONE

=> s 11 ful FULL SEARCH INITIATED 17:45:37 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 2265 TO ITERATE

100.0% PROCESSED 2265 ITERATIONS 766 ANSWERS SEARCH TIME: 00.00.01

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FILE 'CAPLUS' ENTERED AT 17:45:42 ON 17 JUL 2008
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FILE COVERS 1907 - 17 Jul 2008 VOL 149 ISS 3 FILE LAST UPDATED: 16 Jul 2008 (20080716/ED)

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=> s 13 L4 15 L3

=> d bib abs 1-15

- ANSWER 1 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- 2007:1420376 CAPLUS AN
- DN 148:24453
- Treatment of gastrointestinal disorders with CGRP-antagonists
- Doods, Henri; Arndt, Kirsten; Bouyssou, Thierry; Mueller, Stephan Georg; Rudolf, Klaus; Schaenzle, Gerhard
- Boehringer Ingelheim International G.m.b.H., Germany; Boehringer Ingelheim PA Pharma G.m.b.H. & Co. K.-G. SO PCT Int. Appl., 35pp.
- CODEN: PIXXD2
- Patent T.A English

F.	AN.		2																
		PA:	TENT :	NO.			KIN	D	DATE			APPL	ICAT:	ION :	NO.		D	ATE	
								-											
Ρ	I	WO	2007	1412	85		A1		2007	1213		WO 2	007-	EP55	543		2	00706	606
			W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,	CA,
				CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,	ES,	FI,
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				TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW					
			RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
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		US	20080139591				A1		2008	0612		US 2	007-	7600	54		2	0070	808
		US	20080139537				A1		2008	0612		US 2	007-	7600	57		2	00706	808
P	RAI	EP	2006-11787				A		2006	0608									

The invention relates to a method for preventing and treating visceral pain and gastrointestinal disorders such as functional bowel disorders and inflammatory bowel diseases through the use of effective amts. of a compound acting as CGRP antagonist. Twenty eight compds. are calimed (no data). THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE.CNT 4 ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN T. 4 AN 2007:1176398 CAPLUS

- DN 147:455648
- New crystalline compounds of CGRP antagonists
- Ries, Üwe; Sproll, Sonja; Werthmann, Ulrike; Zopf, Andreas; Huchler, TN
- Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G., Germany PA
- SO Ger. Offen., 68pp. CODEN: GWXXBX
- DT Patent
- LA German
- FAN CNT 1

L PHI	PATENT NO.	KIND	DATE	APPLICATION NO.	
PI	DE 1020060178 WO 2007118819	327 A1 9 A2	20071018 20071025	DE 2006-10200601 WO 2007-EP53488	7827 20060413
	WO 2007118819 W: AE, A		20080529 AT, AU, AZ,	BA, BB, BG, BH, BR,	BW, BY, BZ, CA,
				DK, DM, DZ, EC, EE, HU, ID, IL, IN, IS,	
				LR, LS, LT, LU, LY, NI, NO, NZ, OM, PG,	
	RS, E		SE, SG, SK,	SL, SM, SV, SY, TJ,	
	RW: AT, E	BE, BG, CH,	CY, CZ, DE,	DK, EE, ES, FI, FR,	
	BJ, (	CF, CG, CI,	CM, GA, GN,	NL, PL, PT, RO, SE, GQ, GW, ML, MR, NE,	SN, TD, TG, BW,
				SD, SL, SZ, TZ, UG, AP, EA, EP, OA	ZM, ZW, AM, AZ,
DDAT	US 2008008600		20080410	US 2007-734520	20070412

PRAI DE 2006-102006017827 A 20060413

AB The invention concerns novel crystalline compds. of CGRP antagonists that are prepared as salts of hydrochloric acid, hydrobromic acid, sulfuric acid, phosphoric acid, benzene sulfonic acid, p-toluene sulfonic acid, maleic acid, succinic acid, fumaric acid, D-(-)-tartaric acid, L-(+)-tartaric acid, naphthalene 2-sulfonic acid and naphthalene-1,5-disulfonic acid, their polymorph modifications, solvates and hydrates.

- L4 ANSWER 3 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2007:376170 CAPLUS
- DN 146:402012
- Preparation of benzodiazepinones, quinolones, quinazolones, and related compounds as calcitonin gene-related peptide (CGRP) receptor antagonists
- Mueller, Stephan Georg; Rudolf, Klaus; Lustenberger, Philipp; Schaenzle, Gerhard; Stenkamp, Dirk; Doods, Henri; Arndt, Kirsten
- PA Boehringer Ingelheim International G.m.b.H., Germany; Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G.
- PCT Int. Appl., 142pp. SO CODEN: PIXXD2
- Patent DT
- LA German
- FAN CNT 1

	PA:	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE		
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PI	WO	2007	0365	33		A2		2007	0405		WO 2	006-	EP66	789		2	0060	927	
	WO	2007	0365	33		A3		2007	0607										
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			MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RS,	
	RU, SC, SD					SE,	SG,	SK,	SL,	SM,	SV,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	
			UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW								

RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AP, EA, EP, OA EP 1770087 A1 20070404 EP 2005-21282 20050929 R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL, BA, HR, MK, YU 20080618 EP 2006-793855 R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR PRAI EP 2005-21282 Α 20050929 WO 2006-EP66789 20060927 MARPAT 146:402012

Title compds. [I; R1 = Q1, etc.; R2 = (substituted) Ph, pyridin-3-y1; R3 = O2, O3; X = N, C; R4 = Ph, pyridinyl; R5 = C(O)OR9; R6 = H, halo, OH, CF3, alkoxy; R7 = H, alkyl, etc.; R8 = a free electron pair if X = N or R8 = H, alkyl if X = C; R9 = H, alkyl, Ph, indanyl, etc.], tautomers, isomers, diastereomers, enantiomers, hydrates, mixts. salts and salt hydrates thereof, in particular salts thereof, which are physiol. compatible with acids or inorg. or organic bases, were prepared Thus, a solution of 4-(2-oxo-1,3,4,5-tetrahydro-1,3-benzodiazepin-3-v1)piperidin-1-carboxylic acid (R)-1-(4-amino-3-chloro-5-trifluoromethylbenzyl)-2-[4-(4ethoxycarbonylphenyl)piperazin-1-yl]-2-oxoethylester (preparation given), TBTU and Et3N in DMF was stirred for 10 min at room temperature followed by stirring with Et 4-piperazin-1-ylbenzoate for 2 h to give 95% I [R1 = Q1; R6 = H; R2 = (4-amino-3-chloro-5-trifluoromethyl)phenyl; R3 = Q2; X = N; R7 = H; R8 = electron pair; R4 = Ph; R5 = 4-ethyloxycarbonyl]. Tested I showed affinity to human CGRP receptors with IC50 ≤10,000 nM.

- L4ANSWER 4 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2007:197982 CAPLUS
- DN 146:274408

os

GI

- Preparation of N-(carbomethoxy)piperidines as CGRP antagonists
- IN Mueller, Stephan Georg; Rudolf, Klaus; Lustenberger, Philipp; Schaenzle, Gerhard; Santagostino, Marco; Stenkamp, Dirk; Arndt, Kirsten; Doods, Henri PΑ Boehringer Ingelheim International G.m.b.H., Germany; Boehringer Ingelheim

Pharma G.m.b.H. & Co. K.-G. PCT Int. Appl., 377pp. CODEN: PIXXD2

DT Patent LA German FAN.CNT 1

KIND DATE APPLICATION NO. PATENT NO. WO 2007020261 20070222 WO 2006-EP65314 20060815 PI A2 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM DE 102005038831 A1 20070222 DE 2005-102005038831 20050817 DE 2005-102005050953 DE 102005050953 A1 20070426 20051025 US 20070049581 A1 20070301 US 2006-462511 20060804 AU 2006281416 20070222 AU 2006-281416 20060815 A1 CA 2618834 20070222 CA 2006-2618834 A1 EP 1917256 20080507 EP 2006-778243 A2 20060815 R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL, BA, HR, MK, RS MX 200801977 Δ 20080325 MX 2008-1977 20080211 KR 2008039990 20080507 KR 2008-706215 20080313 Α PRAI DE 2005-102005038831 A 20050817 DE 2005-102005050953 A 20051025 WO 2006-EP65314 W 20060815 os MARPAT 146:274408 GI

- AB Title compds. I [X = R3-R4; R1 = substituted 3,4-dihydro-IH-quinazolin-2ones, 1,3,4,5-tetrahydro-ZH-benzo-1,3-diazepin-2-ones, etc.; R2 =
  2-chloro-6-methylaniline, 2-chloro-6-(trifluoromethyl)aniline, etc.; R3 =
  substituted piperidines with provisos; R4 = substituted piperidines with
  provisos] and their pharmaceutically acceptable salts and formulations
  were prepared For example, N-(carbomethoxy)piperidine was prepared from
  4-amino-3-chloro-5-trifluoromethylbenzaldehyde in 9-steps. In GCRP
  receptor binding assays, compds. I exhibited IC50 values ≤10000 nM.
- L4 ANSWER 5 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2006:1173484 CAPLUS
- DN 145:489283
- TI N-Acylpiperidines and related compounds as CGRP-antagonists, methods for preparing them, pharmaceutical compositions and their use as pharmaceutical compositions
- IN Mueller, Stephan Georg; Rudolf, Klaus; Lustenberger, Philipp; Stenkamp, Dirk; Santagostino, Marco; Paleari, Fabio; Schaenzle, Gerhard; Arndt, Kirsten; Doods, Henri
- PA Boehringer Ingelheim International GmbH, Germany
- SO U.S. Pat. Appl. Publ., 156pp.
- CODEN: USXXCO DT Patent
- LA English
- LA English

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	WO	2005	0928	80		A1		2005	1006		WO 2	005-	EP30	94		2	0050	323	
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		2005				A2		2005			WO 2	005-	EP41	04		2	0050	418	
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	EP	1770		иш,	ы,	A1		2007	0404		EP 2	005-	2128	3		2	0050	929	
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EP 2005-21283 A 20050929
DE 2004-102004015723 A 20040329
DE 2004-102004019492 A 20040422
MARPAT 145:489283

GI

AR The invention relates to the CGRP-antagonists of general formula I, the tautomers, the isomers, the diastereomers, the enantiomers, the hydrates, mixts, and salts thereof and the hydrates of the salts, particularly the physiol. acceptable salts thereof with inorg, or organic acids or bases, as well as those compds. of general formula I in which one or more hydrogen atoms are replaced by deuterium, pharmaceutical compns. containing these compds., the use thereof and processes for the preparation thereof. Compds. of formula I wherein X is CH2, NH, C1-3 alkyl-N, O and S; R1 is (spiro)substituted piperidine and oxodihydrothienopyrimidinyl; R2 is (un) substituted (un) fused aryl, and (un) substituted (un) fused pyridine; R3 is (un) substituted piperidine, (un) substituted piperazine, and (un) substituted diazepine; R4 is (un) substituted 4- to 7-membered oxycycloalkyl; and their tautomers and pharmaceutically acceptable salts thereof, are claimed. Example compound II was prepared by cyclization of 2-amino-3-methylphenol with CDI; the resulting 4-methyl-3H-benzoxazole-2one underwent bromination to give 6-bromo-4-methyl-3H-benzoxazol-2-one, which underwent coupling with Me 2-acetylaminoacrylate to give Me 2-acetylamino-3-(4-methyl-2-oxo-2,3-dihydrobenzoxazol-6-yl)acrylate, which underwent hydrolysis to give 3-(4-methyl-2-oxo-2,3-dihydrobenzoxazole-6y1)-2-oxopropionic acid, which underwent asym. reduction to give (R)-2-hydroxy-3-(4-methyl-2-oxo-2,3-dihydrobenzoxazol-6-yl)propionic acid, which underwent esterification to give the corresponding Me ester, which reacted with 4-nitrophenyl chloroformate and 3-(piperidin-4-yl)-1,3,4,5tetrahydro-1,3-benzodiazepin-2-one followed by hydrolysis to give (R)-1-carboxy-2-(4-methyl-2-oxo-2,3-dihydrobenzoxazol-6-yl)ethyl 4-(2-oxo-1,3,4,5-tetrahydro-1,3-benzodiazepin-3-y1)piperidine-1-

ΙI

carboxylate, which underwent amidation with 1-(tetrahydropyran-4yl)piperazine to give compound II. All the invention compds. were evaluated for their CGRP binding affinity. The tested compds. exhibited IC50 values ≥ 10 000 nM.

- L4 ANSWER 6 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2006:1005390 CAPLUS
- DN 145:356814
- Preparation of 2-oxo-1,2,4,5-tetrahydro-1,3-benzodiazepin-3-vlpiperidines TI and related compounds as CGRP receptor antagonists
- Mueller, Stephan Georg; Rudolf, Klaus; Lustenberger, Philipp; Stenkamp, Dirk; Santagostino, Marco; Paleari, Fabio; Doods, Henri; Arndt, Kirsten; Schaenzle, Gerhard
- PA Boehringer Ingelheim International G.m.b.H., Germany; Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G.
- SO PCT Int. Appl., 231 pp.
- CODEN: PIXXD2 DT
- Patent
- LA German

	CNT PAT	ENT :	NO.			KIN		DATE					ION :				ATE	
I	WO	2006	1000	09		A1		2006					EP25				0060	
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             IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL,
             BA, HR, MK, YU
     AU 2006226615
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                                                                     20060318
                          A1
     CA 2600909
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                                 20060928
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     EP 1863799
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                                             IN 2007-DN6658
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     MX 200711527
                          Α
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                                             MX 2007-11527
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                                 20071204
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PRAI WO 2005-EP3094
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     WO 2005-EP4104
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     EP 2005-21283
                                20050929
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     DE 2004-102004015723 A
                                20040329
     DE 2004-102004019492 A
                                 20040422
     WO 2006-EP2515
                                20060318
os
    MARPAT 145:356814
GI
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AB Title compds. I [X = CH2, NH, O, etc.; R1 = substituted 2-oxo-1,2,4,5-tetrahydro-1,3-benzodiazepin-3-ylpiperidines, etc.; R2 = 5-methylquinoxalines, 8-methylmidazo[1,2-a]pyridines, etc.; R3 = substituted piperidines, piperazines, etc.; R4 = 4 to 7-membered ocicycloalkyl ring with provisos] and their pharmaceutically acceptable salts and formulations were prepared For example, benzodiazepinylpiperidine II was prepared from 5-amino-m-cresol in 8-steps. In CGRP receptor inhibition assays, compds. I exhibited IC50 values ≤ 10000 nM.

RE.CNI 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L4 ANSWER 7 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2006:1005389 CAPLUS
- DN 145:377393
- TI Preparation of 2-oxo-1,2,4,5-tetrahydro-1,3-benzodiazepin-3-ylpiperidines as CGRP receptor antagonists

- IN Mueller, Stephan Georg; Rudolf, Klaus; Lustenberger, Philipp; Stenkamp, Dirk; Santagostino, Marco; Paleari, Fabio; Dreyer, Alexander; Arndt, Kirsten; Doods, Henri; Schaenzle, Gerhard
- PA Boehringer Ingelheim International G.m.b.H., Germany; Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G.
- SO PCT Int. Appl., 183 pp.

CODEN: PIXXD2

DT Patent LA German

		TENT				KIN	_	DATE			APPL	ICAT	ION :	NO.		D.	ATE	
	WO	2006	1000	26		A1		2006	0928		WO 2	006-	EP25	57		2	0060	321
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,	KR,
			KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,
			MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
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		RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,
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								GN,										
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				ΚZ,	MD,													
	WO	2005				A1		2005			WO 2						0050	
		W:						ΑU,										
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								GR,										
								BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,
				NE,	SN,													
		2006		44		A1		2006			AU 2						0060	
		2600				A1 A1		2006			CA 2						0060 0060	
	EP	1863		-	D.C.			2007			EP 2				O.D.			
		R:						CZ,										
					ы,	ы,	LU,	LV,	PIC,	NL,	PL,	ы,	RU,	SE,	51,	SK,	IK,	DA,
	MO	2007	HR,			А		2007	1010		NO 2	007	2750			2	0070	710
		2007				A		2007			MX 2			6			0070	
		1011				A		2007			CN 2						0070	
		2007				A		2008			KR 2						0070	
ים		2007				A		2007			1/1/ 2	00/-	1242	01			00/1	V22
ΩJ		2003			1570			2003										
		2004			1012	J M		2004										
		RPAT			93	**		2000	0521									
	v.TUT		110.	5,75	,,													

AB Title compds. I [B = substituted Ph, phenols, anilines, etc.; Y = C, N; R3 = cyclopentyl, cyclohexyl, cycloheptyl; R4 = H with provisos] and their pharmaceutically acceptable salts were prepared For example, benzodiazepinylpiperidine II was prepared from 3-trifluoromethylbenzaldehyde in 7-steps. In CGRP receptor inhibition assays, compds. I exhibited IC50 values ≤ 10000 MM.

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L4 ANSWER 8 OF 15 CAPLUS COPYRIGHT 2008 ACS on SIN
- AN 2006:656692 CAPLUS
- DN 145:96491
- TI Use of CGRP antagonists in treatment and prevention of hot flushes in prostate cancer patients
- IN Rudolf, Klaus; Doods, Henri; Mueller, Stephan Georg; Zamponi, Annette; Lustenberger, Philipp; Stenkamp, Dirk; Arndt, Kirsten; Schaenzle, Gerhard; Brickl, Rolf-Stefan
- PA Boehringer Ingelheim International G.m.b.H., Germany; Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G.
- SO PCT Int. Appl., 46 pp.
- CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 1

	PATENT	NO.			KIN	D	DATE			APPL:	ICAT:	ION I	NO.		D	ATE	
						_											
PI	WO 200	50697	54		A1		2006	0706	1	WO 2	005-1	EP13	974		20	0051	223
	WO 200	50697	54		A9		2007	0809									
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,	KR,
		KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,
		MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
		SG,	SK,	SL,	SM,	SY,	ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,
		VN,	YU,	ZA,	ZM,	ZW											
	RW	: AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	IE,
		IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	BJ,
		CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,
		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
		KG,	KZ,	MD,	RU,	ΤJ,	TM,	ΑP,	EA,	EP,	OA						
	DE 102	00406	3755		A1		2006	0720	1	DE 2	004-	1020	0406	3755	20	0041	229
	US 200	50154	921		A1		2006	0713	1	US 2	005-	3014	22		20	0051	213

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CA 2592278
                       A1 20060706 CA 2005-2592278
A1 20070919 EP 2005-843728
                                                                20051223
    EP 1833484
                                                                20051223
        R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
            IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR
                       A1 20071025
                                         US 2007-774995
    US 20070249592
                                                                20070709
PRAI DE 2004-102004063755 A
                             20041229
    US 2005-301422 A1
                             20051213
    WO 2005-EP13974
                        747
                             20051223
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AB The invention discloses a method for treatment or prevention of hot flushes in men who underwent castration, e.g. due to androgen ablation treatment in prostate cancer therapy, comprising administration of an effective amount of a selected CGRP antagonist to the patient, as well as the use of the active compds. for the manufacture of a pharmaceutical composition

intended to be used in this method.

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 9 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN

AN 2006:636811 CAPLUS

DN 145:76714

II Use of selected CGRP antagonists for combating menopausal hot flushes

IN Rudolf, Klaus; Doods, Henri; Mueller, Stephan Georg; Zamponi, Annette; Lustenberger, Philipp; Arndt, Kirsten; Schaenzle, Gerhard; Stenkamp, Dirk; Brickl. Rolf-Skefan

PA Boehringer Ingelheim International GmbH, Germany

O U.S. Pat. Appl. Publ., 21 pp.

CODEN: USXXCO

DT Patent

LA English FAN.CNT 1

		TENT						DATE				ICAT					ATE	
PI	DE CA	2006 1020 2594 2006	0406 097	3752		A1 A1 A1			0713 0713		DE 2 CA 2		1020 2594	0406 097	3752	20 20 20	0051: 0041: 0051:	213 229 223
		W:	AE, CN, GE, KZ, MZ, SG, VN, AT, IS, CF, GM,	AG, CO, GH, LC, NA, SK, YU, BE, IT, CG, KE,	AL, CR, GM, LK, NG, SL, ZA, BG, LT, CI, LS,	AM, CU, HR, LR, NI, SM, ZM, CH, LU, CM, MW,	AT, CZ, HU, LS, NO, SY, ZW CY, LV, GA, MZ,	AU, DE, ID, LT, NZ, TJ, CZ, MC, GN, NA,	AZ, DK, IL, LU, OM, TM, DE, NL, GQ,	BA, DM, IN, LV, PG, TN, DK, PL, GW,	BB, DZ, IS, LY, PH, TR, EE, PT, ML,	BG, EC, JP, MA, PL, TT, ES, RO, MR,	BR, EE, KE, MD, PT, TZ, FI, SE, NE,	BW, EG, KG, MG, RO, UA, FR, SI, SN,	BY, ES, KM, MK, RU, UG, GB, SK, TD,	BZ, FI, KN, MN, SC, US, GR, TR,	CA, GB, KP, MW, SD, UZ, HU, BF, BW,	CH, GD, KR, MX, SE, VC, IE, BJ, GH,
	EP	1833 R:	483 AT,	BE,	BG,		CY,	2007 CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	
PRAI		2004 2005	-102	0040	6375			2004	1229	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR	

AB The invention discloses the use of selected CGRP antagonists, the physiol. acceptable salts thereof or the hydrates or the hydrates of the salts thereof for combating menopausal hot flushes. A variety of formations are included.

L4 ANSWER 10 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN

AN 2006:636805 CAPLUS

- DN 145:96481
- TI Use of selected CGRP antagonists in combination with other antimigraine drugs for the treatment of migraine
- IN Rudolf, Klaus; Doods, Henri; Mueller, Stephan Georg, Zamponi, Annette; Lustenberger, Philipp; Arndt, Kirsten; Schaenzle, Gerhard; Stenkamp, Dirk; Brickl, Rolf-Stefan
- PA Boehringer Ingelheim International GmbH, Germany
- SO U.S. Pat. Appl. Publ., 22 pp.
- CODEN: USXXCO
- DT Patent
- LA English FAN.CNT 1

PAIN.	PA:	ENT I						DATE				ICAT					ATE	
PI		2006						2006	0629			2005-					0051	216
	DE	1020	0406	3753		A1		2006	0713		DE 2	2004-	1020	0406	3753	2	0041	229
	CA	2594	096			A1		2006	0713		CA 2	2005-	2594	096		2	0051	223
	WO	2006	0724	13		A1		2006	0713		WO 2	2005-	EP13	964		2	0051	223
		W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,	KR,
			KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,
			MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
			SG,	SK,	SL,	SM,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,
			VN,	YU,	ZA,	ZM,	ZW											
		RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,
			IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	BJ,
			CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,
			GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
			KG,	KZ,	MD,	RU,	TJ,	TM										
	EP	1833	478			A1		2007	0919		EP 2	2005-	8232	28		2	0051	223
		R:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
			IS,	IT,	LI,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR	
	US	2008	0103	134		A1		2008	0501		US 2	2007-	9626	33		2	0071	221
PRAI	DE	2004	-102	0040	6375	3 A		2004	1229									
	US	2005	-275	169		B1		2005	1216									
	WO 2005-EP13964 V							2005	1223									
AB	The	inv	enti	on d	iscl	oses	ar	roce	ss f	or t	he t	reat	ment	or	prev	enti	on o	£

AB The invention discloses a process for the treatment or prevention of indications which are selected from among the group comprising headaches, migraine and cluster headaches, the process comprising the joint administration of a therapeutically effective amount of a selected CGRP antagonist (A), a physiol. acceptable salt thereof or a hydrate of the salt and a therapeutically effective amount of a second or third active anti-migraine medicament (B), particularly sumatriptan, zolmitriptan, or dihydroergotamine, or a physiol. acceptable salt thereof, as well as the corresponding pharmaceutical compns. and the preparation thereof. A variety of formulations are included.

- L4 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2005:1154551 CAPLUS
- DN 143:422350
- TI Preparation of 1,3-dihydro-3-(4-piperidiny1)-2H-imidazo[4,5-c]quinolin-2ones and related compounds as cdrp antagonists
- IN Rudolf, Klaus; Mueller, Stephan Georg; Lustenberger, Philipp; Stenkamp, Dirk; Schaenzle, Gerhard; Arndt, Kirsten; Doods, Henri
- PA Boehringer Ingelheim International GmbH, Germany; Boehringer Ingelheim Pharma Gmbh & Co. KG
- SO PCT Int. Appl., 185 pp.
- CODEN: PIXXD2
- DT Patent
- LA German

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FAN.CNT 1
     PATENT NO. KIND DATE APPLICATION NO. DATE
     WO 2005100352 A1 20051027 WO 2005—EP3759 20050409
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
              CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
              GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA,
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              ZM. ZW
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     DE 102004018796 A1
                                   20051103 DE 2004-102004018796 20040415
     CA 2563386 A1 20051027 CA 2005-2563386 20050409 EP 1737860 A1 20070103 EP 2005-729383 20050409
          R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BA, HR, YU
IS, II, LI, LI, LU, MC, NL, PL, FT, RO, SE, SI, SK, TR, BA, HR, JP 2007532601 T 20071115 JP 2007-507727 20050405 US 20050250763 A1 20051110 US 2005-107052 20050415 US 2004-102004018796 A 20040415 US 2004-050948P P 200400511 NO 2005-EP3759 W 20050409 OS MARPAT 143:422350
GI
* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
     Title compds. I [A = substituted Ph, i.e., CF3, NH2, Cl, etc.; X = O, CH2,
AR
     NH; R1 = 1,3-dihydro-2H-imidazo[4,5-c]quinolin-2-onyl,
     1,3-dihydro-2H-benzimidazol-2-one, etc.; NR2R3 = 1,4'-bipiperidiny1,
     1-methyl-4-(4-piperidinyl)piperazinyl, 1-(1-methyl-4-
     piperidinyl)piperazinyl, etc.] and their pharmaceutically acceptable salts
     and formulations were prepared For example, coupling of 1,4'-bipiperidine
     and acid II afforded imidazo[4,5-c]quinolin-2-one III in 76% vield. In
     human cgrp receptor assays, compds. I exhibited IC50 values ≤ 1000
     nM.
RE.CNT 5
               THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
               ALL CITATIONS AVAILABLE IN THE RE FORMAT
T. 4
     ANSWER 12 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
AN 2005:1152762 CAPLUS
DN
    143:440448
     Preparation of 3-piperidin-4-y1-1,3,4,5-tetrahydro-1,3-benzdiazepin-2-ones
TI
     and related compounds as CGRP antagonists
     Mueller, Stephan Georg; Rudolf, Klaus; Lustenberger, Philipp; Stenkamp,
     Dirk; Arndt, Kirsten; Doods, Henri; Schaenzle, Gerhard
     Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G., Germany
PA
     Ger. Offen., 51 pp.
     CODEN: GWXXBX
     Patent
DT
LA German
FAN.CNT 1
     PATENT NO. KIND DATE APPLICATION NO. DATE
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20040415
PΤ
    DE 102004018795
                        A1 20051027 DE 2004-102004018795
     CA 2562526
                          A1
                               20051027 CA 2005-2562526
20051027 WO 2005-EP3741
     WO 2005100343
                          A1
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             NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL,
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             ZM. ZW
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             AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
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                              20070103 EP 2005-731650
     EP 1737842
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                                           JP 2007-507723
     JP 2007532600
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                               20071115
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     US 20050282857
                          A1
                                20051222
                                            US 2005-107195
                                                                     20050415
     US 20070238715
                                            US 2007-688123
                                                                    20070319
                          A1
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PRAI DE 2004-102004018795 A
                                20040415
     US 2004-570005P P
                                20040511
    WO 2005-EP3741
                          W
                                20050409
                         B1 20050415
     US 2005-107195
    MARPAT 143:440448
OS
GT
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- \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT \*
- AB Title compds. I [A = substituted Ph, i.e., CF3, NH2, Cl, etc.; X = 0, CH2, NH; Rl = 3,4-dihydro-2(1H)-quinzolinonyl, 1,34,5-tetrahydro-2H-benzo-1,3-diazepin-2-onyl; NRZR3 = 1,4'-bipiperidinyl, 1-methyl-4-(4-piperidinyl)piperazinyl, 1-(1-methyl-4-piperidinyl)piperazinyl, etc.] and their pharmaceutically acceptable salts and formulations were prepared For example, coupling of 4-(2-piperidin-1-yl-ethyl)piperidine and acid II afforded benzdlazepin-2-one III in 64% yield. In human cgrp receptor assaws, commods. I exhibited IC50 values ≤ 1000 nM.
- L4 ANSWER 13 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2005:1075789 CAPLUS
- DN 143:367334
- TI Preparation of 3-piperidin-4-yl-1,3,4,5-tetrahydro-1,3-benzdiazepin-2-ones as OCGRP receptor antagonists
- IN Mueller, Stephan Georg; Rudolf, Klaus; Lustenberger, Philipp; Stenkamp, Dirk; Dreyer, Alexander; Arndt, Kirsten; Doods, Henri; Schaenzle, Gerhard; Santaqostino, Marco; Paleari, Fabio
- PA Boehringer Ingelheim International G.m.b.H., Germany; Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G.
- SO PCT Int. Appl., 318 pp.
- CODEN: PIXXD2
- DT Patent
- LA German
- FAN.CNT 6

	PATENT	NO.			KIN	D	DATE			APPL	ICAT	I NOI	NO.		D	ATE	
						-											
PI	WO 2005	0928	80		A1		2005	1006		WO 2	005-	EP30	94		2	0050	323
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	WO 2006-EP2557	W	20060321			
OS GI	MARPAT 143:367334					

- \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT \*
- AB Title compds. I [A = O, S; X = O, S; D, E = CH, N with provisos; G = CRa; M = CRb; Q = CRc; Ra, Rb, Rc = H, halo, alkyl, etc.; RI = 5 to 7-membered heterocycle; R2 = H, Ph, pyridinyl, etc.; R3 = H, Ph, pyridinyl, etc.] and their pharmaceutically acceptable salts and formulations were prepared For example, coupling of 1-(1-methylpiperidin-4-yl)piperazine and carboxylic acid II afforded benzdiazepine III in 87% yield. In human OCGRP receptor inhibition assays, compds. I exhibited ICSO values 5 10000 nM.
- RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
  ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L4 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2004:370923 CAPLUS
- DN 140:391302
- TI Preparation of benzo-1,3-diazepin-2-ones and related compounds as CGRP receptor antagonists for the treatment of migraine headaches
- IN Rudolf, Klaus; Mueller, Stephan Georg; Stenkamp, Dirk; Lustenberger, Philipp; Dreyer, Alexander; Bauer, Eckhart; Schindler, Marcus; Arndt, Kirsten: Doods. Henri
- PA Boehringer Ingelheim, Germany
- SO PCT Int. Appl., 254 pp.
- CODEN: PIXXD2
- DT Patent
- LA German
- FAN.CNT 6

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- \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT \*
- AB Title compde. I [A = 0, S, phenylsulfonylimino, etc.; X = 0, S, substituted imino, etc.; Y, Z = alkyl, difluoromethyl, trifluoromethyl, etc.; Rl = 5-7 membered aza, diaza, triaza, etc. heterocycle; R2 = H, phenylmethyl, alkyl, etc.; R3 = H, Ph, pyridinyl, etc.] and their pharmaceutically acceptable salts and formulations were prepared for example, benzo-1,3-diazepin-2-one II was prepared from 1-(3,4-diethylphenyl)lethanone in 8-steps. In human CGRP receptor binding affinity assays, compds. I exhibited ICSO values < 10000 mM. Compds. I are claimed useful for the treatment of migraine headaches.
- RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD
  ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L4 ANSWER 15 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2004:370922 CAPLUS
- DN 140:391301
- TI Preparation of benzo-1,3-diazepin-2-ones and related compounds as CGRP receptor antagonists for the treatment of migraine headaches
- IN Rudolf, Klaus, Mueller, Stephan Georg; Stenkamp, Dirk; Lustenberger, Philipp; Dreyer, Alexander; Bauer, Eckhart; Schindler, Marcus; Kirsten, Arndt; Doods, Henri
- PA Boehringer Ingelheim Pharma G.m.b.H. & Co. K.-G., Germany
- SO PCT Int. Appl., 315 pp.
- CODEN: PIXXD2
- DT Patent
- LA German
- FAN.CNT 1

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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

AB Title compds. I [A = 0, S, phenylsulfonylimino, etc.; X = 0, S, substituted imino, etc.; U = alkyl, alkenyl, alkynyl, etc.; V = Cl, Br, amino, etc.; W = H, halo, difluoromethyl, etc.; R1 = 5-7 membered aza, diaza, triaza, etc. heterocycle; R2 = H, phenylmethyl, alkyl, etc.; R3 = H, Ph, pyridinyl, etc.] and their pharmaceutically acceptable salts and formulations were prepared For example, benzo-l,3-diazepin-2-one II was prepared from 4-amino-3-chloro-5-trifluoromethylbenzoic acid in 9-steps. In human GGRP receptor binding affinity assays, compds. I exhibited IC50 values < 10000 nM. Compds. I are claimed useful for the treatment of migraine headaches.

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L4 ANSWER 15 OF 15 CAPLUS COPYRIGHT 2008 ACS on STN

IT 688019-33-2P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of benzo-1,3-diazepin-2-ones and related compds. as CGRP receptor antagonists for the treatment of migraine headaches)
RN 688019-33-2 CAPUUS

CN 1-Piperidinecarboxylic acid, 4-(1,2,4,5-tetrahydro-2-oxo-3H-1,3benzodiazepin-3-yl)-, (1R)-1-[[4-amino-3-chloro-5-(trifluoromethyl)phenyl]methyl]-2-[4-(4-methyl-1-piperazinyl)-1piperidinyl]-2-oxoethyl ester (CA INDEX NAME)

Absolute stereochemistry.